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journal homepage: www.elsevier.com/locate/apjtbOriginal article <http://dx.doi.org/10.1016/j.apjtb.2015.04.008>Potential of *Hemianax ephippiger* (Odonata-Aeshnidae) nymph as predator of *Fasciola* intermediate host, *Lymnaea natalensis*Aly Younes^{1*}, Hanaa El-Sherif¹, Fathia Gawish², Marwa Mahmoud²¹Department of Entomology, Faculty of Science, Cairo University, Giza, Egypt²Department of Medical Malacology, Theodor Bilharz Research Institute, Giza, Egypt

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ABSTRACT

Objective: To evaluate the predatory capacity of the Odonata, *Hemianax ephippiger* nymph as a biocontrol agent for the freshwater snail *Lymnaea natalensis*, intermediate host of *Fasciola gigantica*.**Methods:** Observations on the searching, attacking and devouring of the snails with a series of laboratory-based predation experiments, whose aims were to determine daily predation rate, differential predation on small-, medium- and large-sized snails were carried out.**Results:** Laboratory evaluation revealed that, the Odonata nymph could kill and consume all three sizes of snails. Searching and handling time of the predator differed depending on snail size and predator vulnerability. The predation rate varied also with respect to snail size and density.**Conclusions:** Our observations suggested that the predator *Hemianax ephippiger* may be a suitable bio-control agent of *Lymnaea natalensis* snail population.

1. Introduction

The liver fluke, *Fasciola gigantica*, is an economically important parasite that infects a wide range of livestock species [1]. The snail *Lymnaea natalensis* Krauss 1848 (*L. natalensis*) functions as an obligatory intermediate host for *Fasciola gigantica* in the old world and thus plays an important role in the epidemiology of *Fasciola* infection [2,3]. Liver flukes can cause huge losses to livestock industries and affect the health of humans where fascioliasis is an important human disease [4–6]. In their general distribution, freshwater pulmonate snails are benthic animals living in the shallower water of lakes, ponds, marshes, rivers, streams and ditches [7,8]. Snail control strategies are considered a priority for the reduction of transmission. Synthetic molluscicides (niclosamide) have been widely used for chemical control [9]; although chemical control only gives a temporary reduction in snail density. The

biological methods, especially those involving the use of indigenous predators, were traditionally perceived as environmentally friendly and have been the foci of research and management of this pest [10]. Predators in nature often include an array of prey types in their diet. Furthermore, in the presence of multiple prey types, they often select certain prey types over others [11]. Predation is a major force affecting species abundance, population dynamics and community structure [12]. Dragonflies are ideal predators of many insect pests and have an important role in biological pests control in various ecosystems. Dragonflies have proved to be potential bio-control agents of mosquitoes and are considered an important predators of various macro-invertebrates [13,14].

Many studies on biological control of freshwater snails using natural enemies have been reported [15–17]. Although some insect predators of snail-host species have been taken into account but other predatory insects like Odonata and Dytiscidae require further study [18]. In view of this, the present study was aimed at evaluating the predation potential of the dragonfly *Hemianax ephippiger* Burmeister 1839 (*H. ephippiger*) nymph of the *Fasciola* snail intermediate host, *L. natalensis*. The searching behavior of the predator towards the different snail

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sizes was also measured. The results of the present study will provide a primary basis for assessment of this predator as biological resource against freshwater snails.

2. Materials and methods

2.1. Collection of *Odonata* nymphs

H. ephippiger nymphs were collected from ponds and lakes in the Abou-Roash area, Giza Governorate, Egypt. They were kept in a glass aquarium (50, 30 and 20 cm in length, width and height), respectively. Collected nymphs were reared in an aquarium in the laboratory and fed daily to satiation on different sizes of *L. natalensis*. Nymphs were starved for a period of 24 h before they were used in experiments. Fully grown nymphs (the last two instars) with sizes ranging from 3.2 to 4.5 cm in length were used in the experiments.

2.2. Collection of snails

The experimental snails, *L. natalensis* were collected from lakes and ponds in the Abou-Roash area, Giza Governorate, Egypt. They were kept in glass aquaria (50, 30 and 20 cm), filled with pond water up to 15 cm of height for a period of one week prior to the start of the experiment. The snails were provided with fresh lettuce leaves as a basic food, dried lettuce is provided when the green was not available. Fish food (Tetramin®) and blue green algae (*Nostoc muscorum*) were used as an additional food source for newly hatched and juvenile snails. Only laboratory-bred snails were used as preys for the nymphs used in experiments. Additionally, some water plants (*Ceratophyllum demersum* and *Elodea* sp.) were placed in the aquarium to simulate natural conditions. Small-, medium- and large-sized snails measuring 2–5, 6–9 and 10–13 mm in shell height, respectively, were used in the experiments.

2.3. Experimental methods

Ten glass aquaria, 5 L in total volume, containing 3 L of pond water were used in each experiment. Among these, the experimental group was comprised of five glass aquaria, each containing a predator and experimental snails. The remaining five glass aquaria constituted the control with only snails. The aquaria were covered with nylon net to prevent snail escape. Snails that may leave the water and sit on the aquarium wall were not considered and deleted from the count. The snails were allowed to acclimatize for 1 h before introducing the predator. Snails and *Odonata* nymphs were used only once in the experiments. All experiments were carried out at constant temperature of $(25 \pm 2)^\circ\text{C}$, 60%–70% relative humidity. Fluorescent tubes (10 cm long, 32 watt, were placed 100 cm above the tanks to provide a photo period of L12: D12.

2.4. Searching and handling time

Predator and prey behaviors were observed during a continuous 60-min period. Foraging behaviors (searching and handling prey) for both starved and satiated predators were quantified. Handling time per prey was calculated as the total time taken to manipulate a single prey item, from encounter to the end of consumption. Encounters between predators and prey,

and the outcomes of the encounters, were also quantified. The encounter rate was calculated as the total number of encounters divided by predator search time (no./min). Encounters with prey could result in attacking, pre-capture, avoidance or consumption of prey. Each trial involved introducing an individual *Odonata* nymph into the experimental glass aquaria filled with clear pond water (to facilitate observation) and containing 10 live snails of one of the three different snail's sizes.

2.5. Effect of prey density on the consumption and predation rate

For each *H. ephippiger* nymph, *L. natalensis* snails (small, medium or large) were supplied at densities of 5, 10, 15, 20 and 25 snails. Predators were allowed to prey for a period of 24 h. Five replicates for each prey density were performed to determine the mean number of prey consumed/day and subsequently the predation rate.

2.6. Data analysis

Data considering searching, handling times, foraging behavior, prey consumed and predation rate were expressed as mean \pm SE. Comparison between three or more different groups was analyzed using One-way ANOVA followed by Bonferroni multiple comparison test for least significant difference. The correlation between prey density and predation rate was determined. Data were analyzed using GraphPad InStat software (version 3.1, GraphPad InStat., California, USA).

3. Results

3.1. Searching and handling times

The *Odonata H. ephippiger* nymphs showed clear differences in searching and handling times towards the three-prey sizes (Table 1). Data obtained show that, the nymph required more time in searching for the small and medium snails as compared to the large snails. The maximum searching time (21.40 ± 1.90) min was obtained by the predator nymph towards small snail when predator nymphs were satiated, whereas the minimum searching time (6.00 ± 0.71) min was obtained with the large snail when the predator nymphs were starved. Significant differences ($P < 0.05$) were obtained in the handling time of the predator nymph towards the three sizes of snails. Handling time of the starved predators towards the snails was (6.80 ± 0.86),

Table 1

Searching and handling times of the predator, *H. ephippiger* nymph towards *L. natalensis* snails.

Behavior	Adult condition	Prey snail size		
		Small	Medium	Large
Searching time (min)	Starved	15.00 ± 1.73^a	15.40 ± 0.93^a	6.00 ± 0.71^b
	Satiated	$21.40 \pm 1.90^{a*}$	18.80 ± 1.20^a	$12.40 \pm 1.20^{b*}$
	P-value	0.037	0.127	0.003
Handling time (min)	Starved	6.80 ± 0.86^a	7.80 ± 1.20^a	12.00 ± 0.71^b
	Satiated	8.80 ± 1.10^a	$11.20 \pm 0.60^{a*}$	$15.40 \pm 0.40^{b*}$
	P-value	0.183	0.018	0.005

Means in the same row, followed by different letters are significantly different ($P < 0.05$); *: Significant at $P < 0.05$ compared with starved value.

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